

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶ : H01J	A2	(11) International Publication Number	WO 99/67801
1101		(43) International Publication Date:	29 December 1999 (29.12.99)

CH

- (21) International Application Number: PCT/US99/13965
- (22) International Filing Date: 21 June 1999 (21.06.99)
- (71) Applicant (for all designated States except US): IONWERKS

22 June 1998 (22.06.98)

- (71) Applicant (for all designated States except US): IONWERKS [US/US]; Suite 255, 2472 Bolsover, Houston, TX 77005 (US).
- (72) Inventor; and

(30) Priority Data:

1328/98

- (75) Inventor/Applicant (for US only): GONIN, Marc [CH/US]; 2215 Addison Road, Houston, TX 77030 (US).
- (74) Agent: PRICE, Ricardo, A.; Fulbright & Jaworski, L.L.P., Suite 5100, 1301 McKinney, Houston, TX 77010 (US).

(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

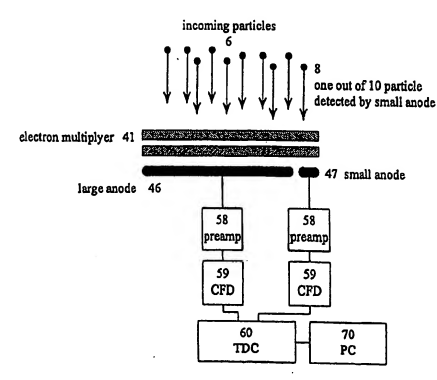
Published

Without international search report and to be republished upon receipt of that report.

(54) Title: A MULTI-ANODE DETECTOR WITH INCREASED DYNAMIC RANGE FOR TIME-OF-FLIGHT MASS SPECTROMETERS WITH COUNTING DATA ACQUISITION

(57) Abstract

new detection scheme for time-of-flight mass spectrometers is disclosed. This detection scheme allows extending the dynamic range spectrometers operating with a counting technique (TDC). The extended dynamic range is achieved by constructing a multiple anode detector wherein the individual anodes detect different fractions of the incoming particles. Different anode fractions are achieved by varying the size, physical location, and electrical/magnetic fields of the various anodes. An anode with a small anode fraction avoids saturation and allows an ion detector to render an accurate count of ions even for abundant species.



FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	Fl	Finland	LT	Lithuania	SK	Slovakia
ΑT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑÜ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IB	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	1L	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Vict Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
СН	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	211	Zimbaowe
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ.	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	u	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SB	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

APPLICATION FOR PATENT

TITLE:

A MULTI-ANODE DETECTOR WITH INCREASED DYNAMIC RANGE FOR TIME-OF-FLIGHT MASS SPECTROMETERS WITH COUNTING DATA ACQUISITION

SPECIFICATION BACKGROUND OF THE INVENTION

1. Field of the Invention

5

The present invention is useful in time-of-flight mass spectrometry (TOFMS), a method for qualitative and quantitative chemical analysis. Many TOFMS work with counting techniques, in which case the dynamic range of the analysis is strongly limited by the measuring time and the cycle repetition rate. This invention describes a detection method to increase the dynamic range of elemental-, isotopic-, or molecular analysis with counting techniques.

10

2. Description of the Prior Art

5

10

15

20

25

Definition of Terms

Anode: The part of a particle detector, which receives the electrons from the electron multiplier.

Anode Fraction: The fraction of the total amount of particles, which is detected by a specific anode.

Single Signal: The signal pulse produced by a detector when a single particle hits the detector. A counting electronics counts the single signals and their arrival.

Signal: A superposition of single signals, caused by particles of one specie hitting the detector within a very short time.

Description

Time-of-flight mass spectrometers (TOFMS, see Figure 1) allow the acquisition of wide-range mass spectra at high speeds because all masses are recorded simultaneously. Most TOFMS work in a cyclic mode. In each cycle, a certain number of particles (up to several thousand) are extracted and traverse a flight section towards a detector. Each particle's time-of-flight is recorded to deliver information about its mass. Thus, in each cycle, a complete time spectrum is recorded and added to a histogram. The repetition rate of this cycle is commonly in the range of 1 to 50 kHz.

If several particles of one specie are extracted in one cycle, these particles will arrive at the detector within a very short time period (as short as 1 nanosecond). When using an analog detection scheme (transient recorder, oscilloscope) this does not cause a problem because these detection schemes deliver a signal which is proportional to the number of particles arriving within a certain sampling time. However, when a counting detection scheme is used (time-to-digital converter, TDC), the electronics cannot distinguish two or more particles of the same specie arriving simultaneously at the detector. Additionally, most TDCs have dead times (typically 20 nanoseconds), which prevent the detection of more than one particle or each mass in one extraction cycle.

For example, when analyzing an air sample with 12 particles per cycle, there will be approximately ten nitrogen molecules (80% N_2 in air, mass = 28 amu) per extraction cycle. These ten N₂ particles will hit the detector within 2 nanoseconds (in a TOFMS of good resolving power). Even a fast TDC with only 0.5 nanoseconds timing resolution and no deadtime will not be able to detect all these particles because only one signal can be recorded each 0.5 nanoseconds. The detection system gets saturated at this intense peak. Figure 2 shows these ten particles 5 of mass 28 amu impinging a detector of prior art. The TDC will register only the first of all these ten particles. Therefore peaks for abundant specie (N₂ and 0₂) are artificially small and are recorded too early because only the first particle is registered. This effect is termed "saturation." Figure 9 shows the effects of saturation on the spectrum peaks for N₂ and O₂. To give a better overview, three different scalings of the same spectrum are shown. The abundance should be 78% N₂, 21% O₂ and 1% Ar. As shown in Figure 9, the N₂ peak and the 0₂ peak are much too small compared to the Ar peak which is not saturated (top and bottom panel). Saturation is so strong that there are virtually no counts during the dead time of approximately 20 nanoseconds registered (middle panel).

5

10

15

20

25

In an attempt to prevent saturation, some prior art detectors use multiple anodes. An individual TDC channel records each anode. Figure 3 shows a prior art detector with four anodes of equal size. This allows the identification of four times larger intensities compared with a single anode detector. However, even with four anodes, the detection of the ten N_2 particles leads to saturation because there are more than two particles per anode on average 6 and 7.

With more anodes, saturation could in principle be avoided, but as each anode requires its own TDC channel, this solution becomes complex and expensive.

SUMMARY OF THE INVENTION

Instead of using multiple equal sized anodes, the present invention uses multiple anodes wherein each anode has a different anode fraction. By reducing anode fraction, saturation can be eliminated. One method for achieving a different anode fraction is through use of anodes of different sizes as shown in Figure 4 at 46 and 47. The example in Figure 4 uses two unequal size anodes with a size ratio of approximately 1:9. As a result, the small anode only detects one particle 8 per cycle, just on the edge of saturation for N₂. Less abundant particles like Ar (1% abundance in air = 0.12 particles per cycle) are primarily detected and evaluated from the big anode which gives low statistical errors. Thus, with 2 anodes of unequal size it is possible to increase the dynamic range by a factor of ten or more. A prior art detector with equal sized anodes would require ten anodes to obtain the same improvement. It should be apparent that the dynamic range can be increased either by decreasing the anode fraction of the small anode or by adding additional anodes with even lower anode fractions. It is also possible to achieve differing fractions and to make such fractions adjustable by applying electric fields to influence the paths of incoming ions as shown in Figures 6 and 7.

20

25

5

10

15

BRIEF DESCRIPTION OF THE DRAWINGS

A better understanding of the present invention can be obtained when the following detailed description of the preferred embodiment is considered in conjunction with the following drawings in which:

Figure 1 is a schematic diagram showing a time-of-flight mass spectrometer to which the invention can be advantageously applied;

Figure 2 is a schematic diagram showing a single anode detector of the prior art;

Figure 3 is a schematic diagram showing a multiple anode detector of the prior art;

Figure 4 is a schematic diagram showing a detector with multiple, unequalsized anodes in accordance with the present invention;

Figure 5 is a graph showing a generic spectrum including an 80% component and a 1 ppm component to depict the saturation effects suffered by prior art detectors and the spectrums generated by a detector of the present invention;

5

10

15

20

25

٩

Figure 6 is a schematic diagram showing an alternate embodiment of the detector of the present invention with two large anodes and one small anode;

Figure 7 is a schematic diagram showing numerical simulations of two electron paths of the detector of Figure 6 achieved by varying the electrical field within the detector;

Figure 8 is a flowchart showing a method for evaluating the spectra taken with a 2-anode detector with unequal-sized anodes; and

Figure 9 is a graph showing a sample spectrum of air with two saturated peaks $(N_2 \text{ and } 0_2)$.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

Referring now to Figure 1, a typical TOFMS is shown. In the depicted TOFMS, gaseous particles are ionized and accelerated into a flight tube from an extraction chamber 20. Some TOFMS, such as the one illustrated, use reflectors to increase the apparent length of the flight tube and, hence, the resolution of the TOFMS. At the detector of the TOFMS 40, ions 6 impinge an electron multiplier 41 causing an emission of electrons. Anodes 44 detect the electrons from the electron multiplier 41 and the signal is then processed through a preamplifier 58, a CFD (constant fraction discriminator) 59, and a TDC 60. A histogram that reflects the composition of the sample is generated either in the TDC 60 or in a digital computer (PC) 70 connected to the TDC 60.

One preferred embodiment of the present invention is shown in Figure 4. In this embodiment, unequal-sized anodes 46 and 47 are used in the detector. The detection fraction of the small anode is small enough so that on average it detects only

one particle 8 out of the ten incoming particles 6 of the specie. The embodiment shown in Figure 4 uses two unequal size anodes with a size ratio of approximately 1:9. As a result, the small anode only detects one particle 8 per cycle, just on the edge of saturation for N_2 . Less abundant particles like Ar (1% abundance in air = 0.12 particles per cycle) are primarily detected and evaluated from the big anode which gives low statistical errors. Thus, with 2 anodes of unequal size it is possible to increase the dynamic range by a factor of ten or more.

5

10

15

20

25

Figure 5 shows the results achieved by using the detector of Figure 4. The top graph of Figure 5 is in logarithmic scale, while the bottom graph is linear. The spectrum recorded by the large anode is shown as a solid line, while the smaller anode's spectrum is shown as a dashed line. As shown in Figure 5, the large anode becomes saturated in the area of an abundant specie (shown between 2000 and 2060 nanoseconds TOF). However, less abundant specie are recorded accurately by the large anode. Also shown in Figure 5, the anode with the smaller anode fraction records the abundant specie without becoming saturated. Thus, by using anodes with different anode fractions, it becomes possible to create an entire spectrum without saturation effects by evaluating minor species (e.g., 1 ppm) on the large anode and major species on the small anode.

Figure 6 shows an alternate embodiment of the present invention. In this embodiment, the electrical potential applied to the small anode 47 is variable, which gives a method for adjusting the small anode's 47 anode fraction. The lower potential on the small anode 47 is less attractive to the electrons 8 and 9 resulting in detection of a smaller fraction of particles 8 and 9 by the small anode 47. Alternative methods for changing the fractions detected by an anode include the application of magnetic fields and physically constructing the instrument in a way such that the ion beam hits the various anodes with different intensities. In most cases, a mixture of these three methods will be used. For example, the detector shown in Figure 7 varies each anode's anode fraction through a combination of size differences, geometry, and electrical potential.

Figure 8 is a flowchart showing a preferred method for evaluating the spectra taken with a 2-anode detector with unequal anode sizes. The additional procedures are encapsulated in the dashed box. As can be seen from the flowchart, upon creating anode histograms, the histograms are analyzed to detect which spectrum regions reflect large anode saturation. In many cases, certain spectrum regions will theoretically be assumed to be saturated and will be treated as such by the method. However, saturation can also be detected by comparing the large anode histogram to the small anode histogram. The small anode histogram, which is not saturated, will accurately reflect the ratio between counts for various regions. Upon comparing these ratios to the same ratios on the large anode histogram, it becomes apparent which histogram regions are saturated. Saturated regions are evaluated by adding the region's large anode histogram to a weighted small anode histogram for that region. The weighting factor is inversely proportional to the small anode's detection fraction. Unsaturated regions are evaluated by adding the large anode histogram to the unweighted small anode histogram. Finally, the processed regions are merged to form a raw spectrum which is corrected with the instrument's transmission function.

5

10

15

20

25

Although the present invention and its advantages have been described in detail, it should be understood that various changes, substitutions and alterations can be made herein without departing from the spirit and scope of the invention as defined by the appended claims. Moreover, the scope of the present application is not intended to be limited to the particular embodiments of the process, machine, manufacture, composition of matter, means, methods and steps described in the specification. As one of ordinary skill in the art will readily appreciate from the disclosure of the present invention, processes, machines, manufacture, compositions of matter, means, methods, or steps, presently existing or later to be developed that perform substantially the same function or achieve substantially the same result as the corresponding embodiments described herein may be utilized according to the present invention. Accordingly, the appended claims are intended to include within their

scope such processes, machines, manufacture, compositions of matter, means, methods, or steps.

CLAIMS

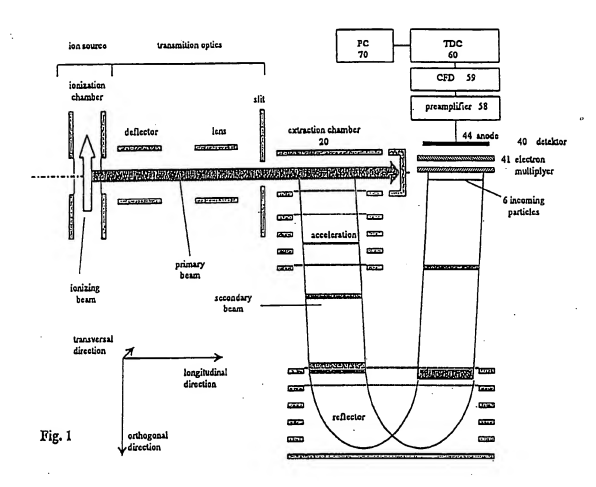
T	- 1		
- 1		aır	m.

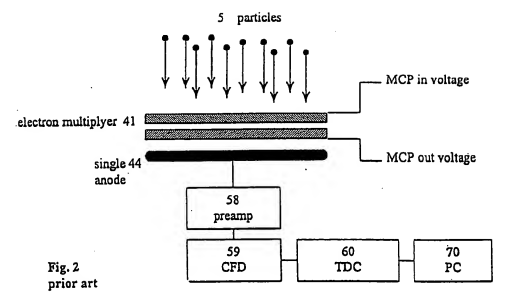
1	1. An ion detector for a time-of-flight mass spectrometer comprising at
2	least two anodes wherein said anodes detect different fractions of incoming ions.
1	2. An ion detector according to claim 1, wherein the size of at least one
2	anode differs from the size of at least one other anode.
1	3. An ion detector according to claim 1, wherein a variable electrical
2	potential on at least one anode modifies incoming ion flight paths such that the anodes
3	detect different fractions of the incoming particles.
1	4. An ion detector according to claim 1, wherein a variable magnetic field
2	in the detector modifies incoming ion flight paths such that the anodes detect different
3	fractions of the incoming particles.
1	5. An ion detector according to claim 1, wherein the ion detector
2	geometry causes the anodes to detect different fractions of the incoming particles.

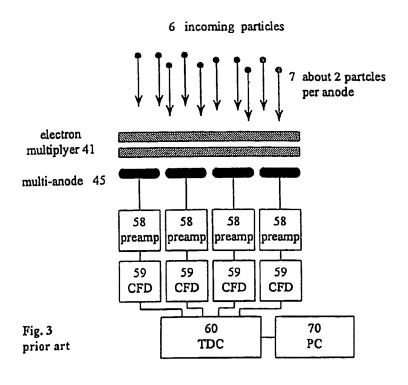
6. A method for creating an ion spectrum in a time-of-flight mass
spectrometer comprising:
(a) recording histograms from at least two anodes wherein said anodes
detect different fractions of incoming ions;
(b) determining which regions of the histogram recorded by at least one
anode that detects a larger fraction of incoming ions are saturated;
(c) creating spectra for saturated regions by applying a weighting factor
to the histogram recorded by the anode that detects a smaller fraction
of incoming ions;
(d) creating spectra for unsaturated regions using unweighted histograms;
and
(e) merging said spectra to form said final ion spectrum.
7. The method of claim 5 wherein said saturation determining step further
comprises treating certain regions as saturated based upon an expected mass
distribution of a sample.
8. The method of claim 5 where said saturation determining step further
comprises comparing the histograms recorded by said anodes on a region by region
basis to create histogram ratios for each region and designating a region as saturated
when its histogram ratio differs substantially from the histogram ratios for other
regions.

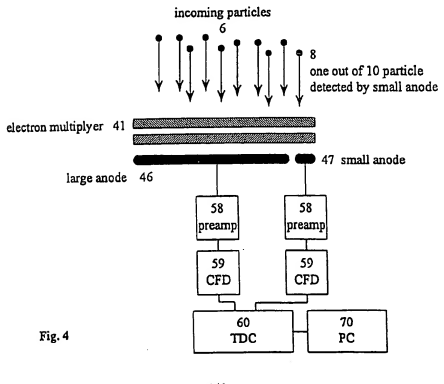
The method of claim 5 wherein said anode frac	tion is determined
theoretically based upon the anode sizes, anode electrical potes	ntials, ion detector
magnetic fields, and ion detector geometry.	

10.	The method of claim 5 wherein said anode fraction is determined
empirically by	comparing histogram peaks for semi-abundant species which are not
so abundant as	to cause saturation on the histogram of the large fraction anode but are
still sufficientl	y abundant so as to register a meaningful result on the small fraction
anode	

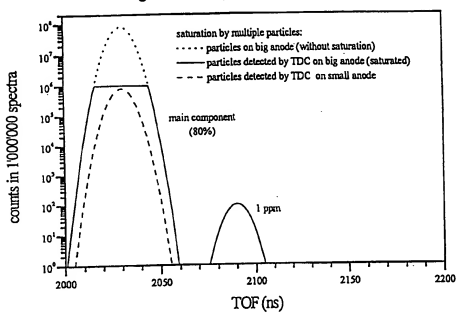








generic mass spectrum in log scale



generic mass spectrum in linear scale

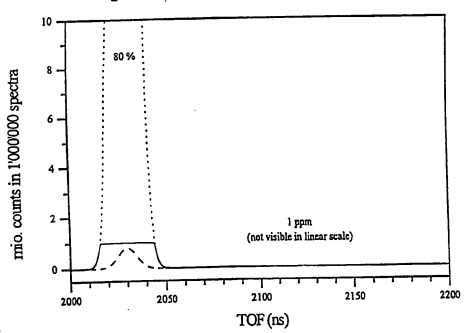
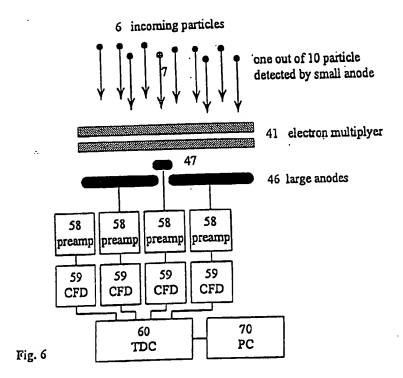
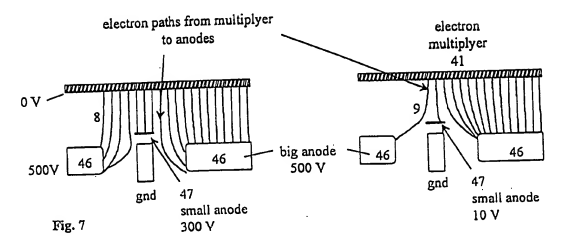
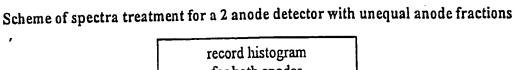
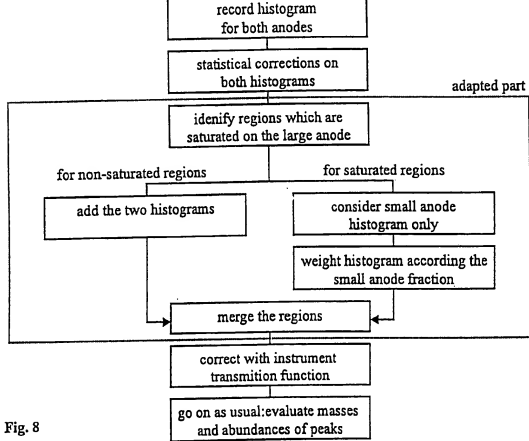


Fig. 5









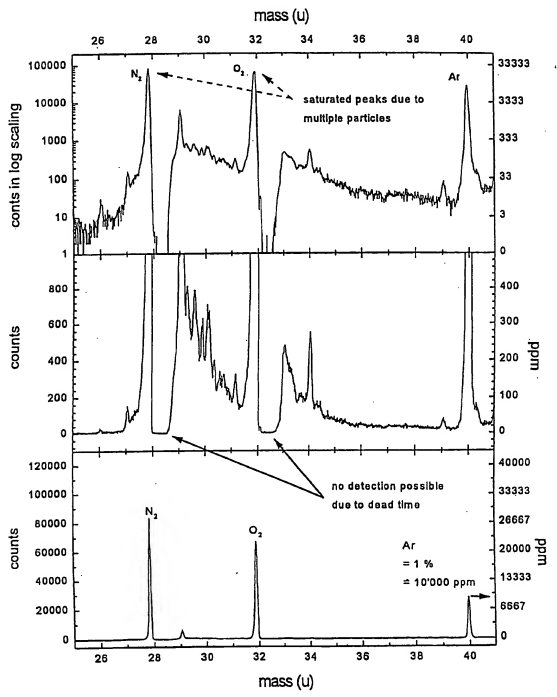


Fig. 9